AMENDED SPECIFICATION (UNDER ARTICLE 34)

MOLDING MATERIAL FOR USE WITH CARBON DIOXIDE REFRIGERANT

TECHNICAL FIELD

The present invention relates to a molding material for use with carbon dioxide refrigerant, and more particularly to a molding material for use with carbon dioxide refrigerant with distinguished resistance to carbon dioxide gas permeation and volumic swelling.

BACKGROUND ART

Freon R-134a (1,1,1,2-tetrafluoroethane) called a new refrigerant is now mainly used in refrigerators, etc., but it is expected that its use would be prohibited in future in view of problems of environmental control, etc. Hydrocarbon gases and carbon dioxide are now regarded as important substitutes for such a refrigerant. Hydrocarbon gases have a considerably high risk of explosion or combustion and the world trend toward the next generation refrigerant is shifting to carbon dioxide.

Carbon dioxide requires a higher pressure than freon and also has higher permeability and solubility into ordinary polymer materials and thus has a higher chance of blister (foaming) generation. Even if there is no blister generation, carbon dioxide gas permeates through the polymer materials, resulting in failure of pressure maintenance or tight sealing.

Even rubber materials have a high carbon dioxide gas permeability in general, and the permeability is considerably high particularly under pressure of 10 atm or higher, resulting in failure of tight sealing of carbon dioxide gas. Furthermore, carbon dioxide is very soluble into polymers,

giving rise to considerable polymer swelling. Thus, no rubber material parts have been actually applied to apparatuses using carbon dioxide.

In case of the conventional freon gas refrigerants, hydrogenated NBR, EPDM, etc. have been used as molding materials for sealing materials, but these rubber materials considerably swell or blister through contact with carbon dioxide as disadvantages.

DISCLOSURE OF THE INVENTION

An object of the present invention is to provide a molding material for use with carbon dioxide <u>refrigerant</u>, which can be used as suitable molding materials for sealing materials, etc. in carbon dioxide <u>refrigerant</u>-contacting apparatuses.

The object of the present invention can be attained by a molding material for use with carbon dioxide refrigerant, which comprises chlorinated polyethylene having a chlorine content of 25-47% by weight. A blend of chlorinated polyethylene with vinyl chloride-based resin can be also used. Furthermore, it is particularly effective for prevention of blister generation to add a specific silane coupling agent thereto.

Chlorinated polyethylene for use in the present invention has a chlorine content of 25-47% by weight, preferably 28-45% by weight. The lower the chlorine content than 25% by weight, the nearer to the property of polyethylene, resulting in loss of rubber elasticity. Sealing materials molded from such chlorinated polyethylene will have poor sealing. With a higher chlorine content than 47% by weight, the low temperature resistance, heat resistance, etc. will be lowered. Actually, commercial products with the above-mentioned range of chlorine content, for example, Elaslene series of Showa Denko products, can be used as such.

Chlorinated polyethylene with such a range of chlorine content can be usually used alone, but in case of improving vulcanization product properties such as tensile strength, elongation, etc. at somewhat the sacrifice of resistances to carbon dioxide gas permeation and volume swelling, a blend of chlorinated polyethylene with vinyl chloride-based resin, usually polyvinyl chloride, can be also used, where a blending proportion of vinyl chloride-based resin is not more than about 50% by weight, preferably about 10 to about 40% by weight, on the basis of the blend.

From the viewpoint of dynamic properties and resistance to gas permeation of chlorinated polyethylene or its blend, about 5 to about 150 parts by weight, preferably about 10 to about 100 parts by weight, of an inorganic filler is added to 100 parts by weight of chlorinated polyethylene or its blend. Reinforcing fillers such as carbon black, silica, etc. can be usually used as an inorganic filler. Furthermore, non-reinforcing fillers such as calcium metasilicate, diatomoceous earth, graphite, mica, calcium carbonate, zinc oxide, etc. can be also used together with the reinforcing fillers.

Furthermore, an organic peroxide is added as a cross-linking agent to the molding material comprising the above-mentioned components. Organic peroxide for use in the present invention includes, for example, 1,1-bis(t-butylperoxy)-3,5,5-trimethylcyclo-hexane, 2,5-dimethylhexane-2,5-dihydroxyperoxide, di-t-butyl peroxide, t-butyl cumyl peroxide, dicumyl peroxide, t-butylperoxyisopropyl carbonate, n-butyl-4,4-di(t-butylperoxy) valerate, α , α -bis(t-butylperoxy)-p-diisopropylbenzene, 1,3-di(t-butylperoxy)-p-diisopropylbenzene, 2,5-dimethyl-2,5-di(t-butylperoxy)-p-diisopropylbenzene, 2,5-dimethyl-2,5-di(t-butylpero

benzene, 2,5-dimethyl-2,5-di(benzoylperoxy)hexane, etc., and is used in a proportion of about 1 to about 10 parts by weight, preferably about 2 to about 8 parts by weight on the basis of 100 parts by weight of chlorinated polyethylene or its blend. In a lower proportion of organic peroxide than about 1 part by weight no satisfactory cross-linking density can be obtained, whereas in a higher proportion than about 10 parts by weight no vulcanization moldings can be obtained due to foaming, or rubber elasticity, elongation or compression set characteristics will be inevitably lowered, even if moldings can be obtained.

To the molding material containing an organic peroxide as a cross-linking agent it is added about 0.1 to about 10 parts by weight, preferably about 1 to about 5 parts by weight, of a polyfunctional unsaturated compound, typically triallyl isocyanurate, triallyl cyanurate, etc. as a cocross-linking agent on the basis of 100 parts by weight of chlorinated polyethylene or its blend. When the cross-linking agent and the cocross-linking agent are used in higher proportions than about 10 parts by weight, respectively, scorching, etc. will take place.

Furthermore, it is effective for prevention of blister generation due to contact with carbon dioxide to add at least 0.2 parts by weight, preferably about 0.5 to about 5 parts by weight, of a specific silane coupling agent to 100 parts by weight of chlorinated polyethylene or its blend with vinyl chloride-based resin.

Such a specific silane coupling agent includes, for example, vinylic silane coupling agents such as vinyltrimethoxysilane, vinyltriethoxysilane, vinyltris(2-methoxyethoxy)silane, vinyltrichlorosilane, etc., epoxy silane coupling agents such as γ -glycidoxypropyltrimethoxysilane, γ -glycidoxypropyltriethoxysilane, γ -glycidoxypropylmethyldiethoxysilane,

etc., and methacryloxysilane coupling agents such as γ -methacryloxypropyltrimethoxysilane, γ -methacryloxypropylmethyldimethoxysilane, γ -methacryloxypropylmethyldimethoxysilane, γ -methacryloxypropylmethyldiethoxysilane, etc.

Chlorinated polyethylene composition comprising the foregoing components can be prepared by further adding thereto, if required, various additives, for example, a processing aid such as stearic acid, palmitic acid, paraffin wax, etc., an acid acceptor such as zinc oxide, magnesium oxide, etc., an antioxidant, a plasticizer, etc. properly, followed by kneading through Intermix, kneader, Banbury mixer, etc. or through open rolls.

Vulcanization of the composition is carried out usually by heating at about 150° to about 200°C for about 3 to about 60 minutes though a vulcanization press, an injection molding machine, a compression molding machine, etc., and, if required, secondary vulcanization is carried out by heating at about 120° to about 200°C for about 1 to about 24 hours.

BEST MODE FOR CARRYING OUT THE INVENTION

The present invention will be described below, referring to Examples.

EXAMPLE 1

| Par | ts by weight | | | |
|---|--------------|--|--|--|
| Chlorinated polyethylene (Elaslene 302NA, | 100 | | | |
| product by Showa Denko K. K.; chlorine content: | | | | |
| 29 wt.%) | | | | |
| Carbon black (Thermax N990, product by Cancarb) | 120 | | | |
| Magnesium oxide (Kyowa Mag #150, product by | 5 | | | |
| Kyowa Kagaku K. K.) | | | | |
| Organic peroxide (Percumyl D. product by NOF | 5 | | | |

Corp.)

Triallyl isocyanurate (Tyke, product by Nihon

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Kasei K. K.)

The foregoing components were kneaded through 10-inch open rolls (roll temperature: 130°C), and the kneading product was press vulcanized at 180°C for 6 minutes to obtain a vulcanization molded circular sheet, 90mm in diameter and 0.5mm thick.

A disc sample, 50mm in diameter, was cut out from the circular sheet and mounted on a high pressure gas permeability tester to determine a CO_2 permeability under high pressure (60 atm). It was found to be $2.7 \times 10^{\cdot 10} \text{cm}^3$ (STP) · cm/cm² · sec · cmHg. Furthermore, a rectangular sample, $20 \text{mm} \times 5 \text{mm}$, was cut out from the circular sheet and exposed to a CO_2 atmosphere under high pressure (60 atm) in a pressurizing apparatus with a window. A volumic swelling rate $\triangle V$ was measured by a cathetometer and found to be 10 vol.%.

EXAMPLE 2

In Example 1, the same amount of Elaslene 406AE, a product by Showa Denko K. K. (chlorine content: 39.5 wt.%) as chlorinated polyethylene and 70 parts by weight of Seest G-S, product by Tokai Carbon Co., Ltd. as carbon black were used. It was found that CO₂ permeability was 4.8×10^{-10} cm³ (STP) · cm/cm² · sec · cmHg and volumic swelling rate \triangle V was 11 vol.%.

EXAMPLE 3

In Example 2, the amount of carbon black was changed to 50 parts by weight and 40 parts by weight of Nipseal ER, product by Nippon Silica K. K. was additionally used as white carbon. It was found that CO₂ permeability was $3.0 \times 10^{-10} \text{cm}^3$ (STP) · cm/cm² · sec · cmHg and volumic

swelling rate $\triangle V$ was 10 vol.%.

EXAMPLE 4

In Example 1, the same amount of Elaslene 452NA, product by Showa Denko K. K. (chlorine content: 44.5 wt.%) was used as chlorinated polyethylene and the amount of carbon black was changed to 90 parts by weight. It was found that CO₂ permeability was $2.2 \times 10^{-10} \text{cm}^3$ (STP) · cm/cm² · sec · cmHg and volumic swelling rate $\triangle V$ was 9 vol.%.

EXAMPLE 5

In Example 1, the amount of chlorinated polyethylene was changed to 70 parts by weight and 30 parts by weight of polyvinyl chloride (ZEST800Z, product by Shin-Daiichi Enbi K. K.) was used as a blend with the chlorinated polyethylene. It was found that CO₂ permeability was 7.6× 10^{-10} cm³ (STP) · cm/cm² · sec · cmHg and volumic swelling rate \triangle V was 16 vol.%.

COMPARATIVE EXAMPLE 1

| | Parts by weight |
|---|-----------------|
| Hydrogenated NBR (Zetpol 1020, product by | 100 |
| Nippon Zeon Co., Ltd.) | |
| Carbon black (Thermax N990) | 80 |
| Magnesium oxide (Kyowa Mag #150) | 5 |
| Organic peroxide (Percumyl D) | 5 |
| Triallyl isocyanurate (Tyke) | 3 |

The foregoing components were subjected to kneading, vulcanization, molding and determination in the same manner as in Example 1, and it was found that CO₂ permeability was $7.3 \times 10^{-9} \text{cm}^3$ (STP) · cm/cm² · sec · cmHg and volumic swelling rate $\triangle V$ was 31 vol.%.

Vulcanization molding products of the foregoing Examples and

Comparative Example were subjected to determination of normal state physical properties according to JIS K-6301. The results are give in the following Table 1.

Table 1

| Example No | <u>Tensile strength (MPa)</u> | Elongation (%) |
|------------|-------------------------------|----------------|
| 1 | 15 | 200 |
| 2 | 16 | 210 |
| 3 | 17 | 190 |
| 4 | 17 | 205 |
| 5 | 22 | 275 |
| Comp. Ex. | 1 18 | 250 |

EXAMPLE 6

| | Parts by weight |
|---|-----------------|
| Chlorinated polyethylene (Elaslene 352NA, | 100 |
| product by showa Denko K. K., chlorine | |
| content: 35 wt.%) | |
| SRF carbon black | 55 |
| Magnesium oxide | 5 |
| Dicumyl peroxide | 4 |
| Triallyl isocyanurate | 5 |
| Vinylic silane coupling agent (vinyl | 2 |
| triethoxysilane) | |

The foregoing components were kneaded through a kneader and open rolls, and the kneading product was press vulcanized at 170°C for 30 minutes and then oven vulcanized (second vulcanization) at 140°C for 10 hours to obtain a vulcanization sheet, 150mm×150mm×2mm.

EXAMPLE 7

In Example 6, the same amount of epoxy silane coupling agent (γ -glycidoxypropyltriethoxysilane) was used in place of the vinylic silane coupling agent.

EXAMPLE 8

In Example 6, the same amount of methacryloxy silane coupling agent (γ -methacryloxypropyltriethyoxysilane) was used in place of the vinylic silane coupling agent.

COMPARATIVE EXAMPLE 2

In Example 6, the amount of dicumyl peroxide was changed to 0.5 parts by weight.

COMPARATIVE EXAMPLE 3

In Example 6, the amount of dicumyl peroxide was changed to 12 parts by weight.

COMPARATIVE EXAMPLE 4

In Example 6, the same amount of amino silane coupling agent (γ -aminopropyltriethoxysilane) was used in place of the vinylic silane coupling agent.

Vulcanization sheets obtained in the foregoing Examples 6 to 8 and Comparative Examples 2 to 4 were subjected to determination and evaluation of the following items. Results are shown in Table 2. In the case of Comparative Example 3, no vulcanization molding could be carried out due to foaming

Normal state physical properties: according to ΠS K-6253 and

JIS K-6251

Compression set: according to JIS K-6262

CO₂ resistance: The sheets were dipped into liquidified CO₂ at

25°C for 24 hours, followed by heating at 150°C for one hour and then by visual inspection of blister generation on the sheet surface

Table 2

| Items of determination | | | | Comp. | . Comp. | |
|------------------------------|------------|------|-------------|-------------|-------------|--|
| and evaluation E | <u>x.6</u> | Ex.7 | <u>Ex.8</u> | <u>Ex.2</u> | <u>Ex.4</u> | |
| [Normal state physical | | | | | | |
| properties] | | | | | | |
| Hardness (durometer A) | 85 | 85 | 84 | 74 | 85 | |
| Tensile strength (MPa) | 20.0 | 20.2 | 20.7 | 24.8 | 17.9 | |
| Elongation (%) | 200 | 210 | 190 | 500 | 350 | |
| [Compression set] | | | | | | |
| 120°C for 70 hours (%) | 20 | 20 | 18 | 67 | 43 | |
| [CO ₂ resistance] | | | | | | |
| Blister generation | None | None | None | None | Occurred | |

INDUSTRIAL APPLICATION

Molding materials for use with carbon dioxide refrigerant, which comprise chlorinated polyethylene, have a CO₂ permeability under 60 atm in the order of 10⁻¹⁰cm³ (STP) · cm/cm² · sec · cmHg or less and a distinguished volumic swelling rate $\triangle V$ of about 10 vol. % or less. When chlorinated polyethylene is used as a blend with vinyl chloride-bassed resin, good vulcanization product properties such as tensile strength: 18 MPa or higher and elongation: 250% or higher can be obtained, though resistances to carbon dioxide gas permeation and volumic swelling may be somewhat sacrificed. Furthermore, when a specific silane coupling agent is added to

chlorinated polyethylene or its blend with vinyl chloride-based resin, blister generation can be effectively prevented.

The present molding materials for use with carbon dioxide refrigerant with the foregoing properties can be used as suitable materials for sealing materials such as packings, gaskets, O-rings, etc. in carbon dioxide refrigerant -contacting apparatuses such as air compressors, refrigerators, supercritical CO₂ extraction (chromatography) apparatuses, etc. using carbon dioxide as a refrigerant, and also as suitable materials for sealing materials for refrigerator oil (polyalkylene glycol, etc.) applicable to the carbon dioxide refrigerant.